## **Chemistry Cheatsheet**

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partly based on the work by L. Hoffmann & D. Vermee

## 1. Basics

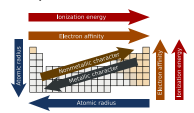
- Energy:  $1eV = 1.602 \cdot 10^{-19} J$ , 1cal = 4.18 J
- Pressure:  $P = \frac{F}{A} = \rho \cdot h \cdot g$

1atm = 760mm Hg = 760torr = 101'325Pa =

- Force:  $F = m \cdot q, \ m = \rho \cdot V$
- Amount of substance:  $1 \text{mol} = 6.022 \cdot 10^{23} \text{(Avogadro)}$
- Length:  $1\text{Å} = 10^{-10} m$
- STP Ideal Gas:  $0^{\circ}C = 273.15K, \ 1atm; \ V_m = 22.41L$
- STP thermodynamics:  $25^{\circ}C = 298K, 1bar, 1mol, 1cal$
- STP electrochemistry:  $25^{\circ}C = 298K$ , 1atm, 1M

- Kinetic energy:  $E_{kin} = \frac{1}{2} \cdot m \cdot v^2$
- Potential energy:  $E_{pot} = m \cdot g \cdot \Delta h$
- electrostatic:  $E_{el} = \frac{\kappa Q_1 Q_2}{d^2}$   $\kappa = \frac{1}{4\pi\epsilon_0}$
- Photon energy:  $E_{\gamma} = h \cdot f = \frac{h \cdot c}{\lambda}$
- De Broglie wavelength:  $\lambda = \frac{h}{m \cdot v}$

- Ionisation energy: The ionization energy is the quantity of energy that an isolated, gaseous atom in the ground electronic state must absorb to discharge an electron, resulting in a cation.
- Electron affinity: Electron affinity is defined as the change in energy (in kJ/mole) of a neutral atom (in the gaseous phase) when an electron is added to the atom to form a negative
- Electronnegativity: Electronegativity is a measure of an atom's ability to attract shared electrons to itself.



### 2. Atoms

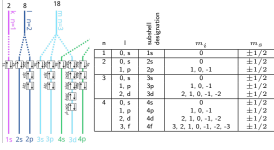
- Atomic number = #protons = #electrons
- mass number = #protons + #neutrons

Heisenbergs uncertainty principle  $\Delta x \cdot \Delta p \geq \frac{h}{4 \cdot \pi}$  Due to duality of electrons (acting like waves and elementary entities at the same time), impossible to exactly describe position and momentum si-

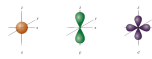
### Effective nuclear charge (approx.): $Z_{eff} = Z - S$

$$Z=\# \mathrm{protons},\, S=\# e^-$$
 on all full shells

In periodic table:  $Z_{eff}$  increases from left to right ightarrow electrons are more attracted and hence atomic radius is smaller, the further right in the periodic table.



- ullet n: principal quantum number o size of orbital
- I: angular quantum number → shape of orbital
- ullet  $m_l$ : magnetic quantum number o orientation of orbital
- m<sub>s</sub>: spin quantum number



Pauli Exclusion: Each electron has unique set of quantum numbers Hund's rule: Every orbital in sublevel is first singly occupied

Energy of Hydrogen  $e^-$ :  $E_n = -\frac{hcR_H}{n^2}$ 

Excitement from shell  $n_1$  to  $n_2$ :  $E_H = hcR_H(\frac{1}{n^2} - \frac{1}{n^2})$ 

### 3. Chemical bondings

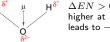
Two atoms share electron pairs octet rule: Atom tries to acquire noble state (2 valence electrons for H and He. 8 valence electrons for all other) exceptions:

- Less / more than 8 VE's on central atom

- Write symbols and connect with single bonds
- Complete octets around non-central atoms
- Place remaining VE around central atom
- Try multiple bonds if central atom does not have octet
- if multiple lewis structures possible: choose most stable according to formal charge

Electrons transfered from atoms with lower EN to atoms with higher EN  $\rightarrow$  cations (+) (smaller radius) and anions (-) (bigger radius). electrostatic attraction.  $\Delta EN > 1.7 \rightarrow$  ionic bonding lattice energy  $\Delta U$ : Energy required to separate ions to infinite distance

cores form grid structure, electron electricity and heat conduction) surrounds atomic cores + + + + +



 $\Delta EN > 0.5$ : polar bonding, density of  $e^$ higher at  $\delta^-$  atom. If molecular structure leads to  $\rightarrow$  dipole moment exists.

Results if new bondings are formed to satisfy octet rule, ignores

### Determine formal charge in Molecule:

- · split all bondings in middle
- ullet atoms formal charge = VE if unpaired  $-e^-$  of that atom after splitting bondings

Bond length:  $\equiv <=<-$ Bond strength:  $- <= <\equiv$ 

**Bond enthalpy** = [broken bonds] - [formed bonds]

#### Most stable molecule:

- least formal charges.
- formal charges necessary: choose smallest effective charge of molecule (sum of formal charges)
- · negative formal charge on electronegative atom

#### 4. Molecular models









structural perspective ball-and-stick

### 4.1 valence shell electron pair repulsion (VSEPR)

Number of	Electron-	Molecular Geometry			
Electron Dense Areas	Pair Geometry	No Lone Pairs	1 Ione Pair	2 Ione Pairs	3 Ione Pairs
2	Linear	• •			
	180°	Linear			
3	Trigonal planar	Trigonal	Bent		
	120°	planar	Don		
	Tetrahedral	Tetrahedral	Trigonal pyramidal	Bent	
5	Trigonal bipyramidal	Trigonal	Seesaw	T-shaped	Linear
	120°/90°	bipyramidal		1-snapcu	Lincus
*	Octahedral 90°	Octahedral	Square pyramidal	Square planar	

#### 5. State of matter

#### 5.1 Intermolecular forces

- 1. Van-der-Waals interactions (weak)
  - (a) Dipol-Dipol

Sum of all dipole moments from polar bonds in molecule. (molecular dipole) ( $\Delta EN > 0.5$ )

### (b) Dispersion

Temporary fluctuations of the electrons can cause an induced dipole. These forces always exist. Force increases with molecule size and also affected by molecular shape.

### 2. Ion-Dipole Interactions (strong)

Very important for solutions. Ions solvated by polar liquid.

### 3. Hydrogen bonding (strong)

One type of dipole-dipole interaction. N, O, F are very electronegativ  $\Rightarrow$  very polar bonds with H.

Ion-Dipole > H-Bonding > Dipole-Dipole  $\approx$  Dispersion > 50kJ/mol  $\sim 25$ kJ/mol  $\sim 50 \text{kJ/mol}$ 

Colligative Properties: Changes depend on amount of solute added, but not which solute.

$$\text{Clausius-Clapeyron} \left\{ \begin{array}{l} \ln(P_{vap}) = -\frac{\Delta H_{vap}}{RT} + C \\ \ln(\frac{P_1}{P_2}) = \frac{\Delta H_{vap}}{R} \left(\frac{1}{T_2} - \frac{1}{T_1}\right) \end{array} \right.$$

 $\Delta T_b = T_b(\text{solution}) - T_b(\text{solvent}) = iK_b m$ 

m = molality of solute

 $K_b = \text{molal bp elevation constant}$ 

i = van't Hoff factor

=1 for non-electrolytes

= Number of ions produced for electrolytes. e.g 2 for NaCl

$$P_{\rm vap}^{\rm sol} = X_{\rm solvent} * P_{\rm vap}^{\rm pure}$$

Raoult's law - Solution is an ideal solution. All intermolecular interactions are identical.

$$\Delta T_f = T_f(solution) - T_f(solvent) = -iK_f m$$

 $m = {\sf molality} \; {\sf of} \; {\sf solute}$ 

 $K_f = \text{molal fp depression constant}$ 

i = van't Hoff factor

### 5.3 Expressions for Solutions

- X Mole fraction =  $\frac{\text{moles solute}}{\text{total moles}}$
- M Molarity =  $\frac{\text{moles solute}}{\text{litres solution}}$
- m Molality =  $\frac{\text{moles solute}}{\text{kg solvent}}$
- Mass\% =  $\frac{\text{mass solute}}{\text{total mass}} \cdot 10^2$ , ppm:  $\cdot 10^6$ , ppb:  $\cdot 10^9$

#### 5.4 Ideal Gas

- Assumptions of IGL
  - Gas molecules don't occupy much of total volume.
  - Gas molecules don't interact.
- Ideal Gas Law: pV = nRT = NkT
- $\bullet \ \ p\left[Pa\right], V\left[m^3\right], n\left[\mathsf{num \ of \ moles}\right], R\left[\tfrac{J}{mol*K}\right], T\left[K\right]$
- Density  $\rho = M \cdot \frac{n}{V} = M \cdot \frac{p}{RT}$

$$p_i = n_i \cdot rac{RT}{V}$$
 total pressure  $= \sum$  of all partial pressures.

Pressure needed to counteract osmotic flow.

$$\Pi = i \left(\frac{n}{V}\right) RT = iMRT$$

#### 6. Thermodynamics

- Open Can echange matter and energy w/ surrounding
- Closed Can echange energy w/ surrounding
- Isolated Nothing can be exchanged

- $\Delta E = E_{\text{final}} E_{\text{initial}}$  $\Delta E > 0$  system gained energy
- $\Delta E < 0$  system lost energy 1st Law of Thermodynamics
- $\Delta E = q + w$

q = heat added to system, w = work done on system

#### 6.3 Enthal

 $\Delta H$  tells us about heat transferred during chemical reaction

- ullet  $\Delta H=q_p$  heat flow at constant P
- $-P\Delta V = \text{pressure-volume work}$
- $\begin{array}{c} \bullet \;\; \Delta H > 0 \Rightarrow \mathrm{endothermic} \\ \Delta H < 0 \Rightarrow \mathrm{exothermic} \end{array}$

#### b.4 Heat Capaci

Heat flow required to raise substance's T by 1 degree  ${}^{\circ}C$  (or K)

$$C_m = \text{molar heat capacity} = \left[\frac{J}{mol \, ^{\circ}\text{C}}\right] = \left[\frac{J}{mol \, K}\right]$$
 $C_s = \text{specific heat capacity} = \left[\frac{J}{a \, ^{\circ}\text{C}}\right] = \left[\frac{J}{a \, K}\right]$ 

$$C_s=$$
 specific heat capacity  $=\left\lfloor \frac{1}{g\ ^{\circ}\mathrm{C}}\right\rfloor =\left\lfloor \frac{1}{g\ K}\right\rfloor$  Ex.  $g=C_{\mathrm{m}}\cdot n\cdot \Delta T$ 

Ex. 
$$q = C_{\mathsf{m}} \cdot n \cdot \Delta$$

Hess's Law:  $\Delta H_{\rm rxn} = \sum \Delta H_i$  e.g Enthalpies of Formation:  $\Delta H_f^{\circ}$ 

#### 6.5 Entropy,

Entropy is a measure of disorder in a system. All spontaneous processes are irreversible. S is a state function.  $\bullet \ \Delta S = \frac{q_{\rm rev}}{r}, \ q_{\rm rev} = {\rm heat \ flow \ for \ reversible \ process}$ 

- $S = k_b \cdot ln(W)$ ,  $k_b = \text{Boltzmann's constant}$ , W = num of
- $b = k_b \cdot tin(w)$ ,  $k_b = Boltzmann s constant$ , w = num c microstates

### $\Delta S > 0$ : increasing microstates

e.g increasing V, increasing T, increasing n, increasing complexity of molecules, melting solids, vaporizing liquids

### 6.6 2nd Law of Thermodynamic

Entropy of the universe increases for any spontaneous process.

$$\Delta S_{
m univ} = \Delta S_{
m sys} + \Delta S_{
m surr} > 0$$
 (spontaneous, irreversible)  
 $\Delta S_{
m univ} = \Delta S_{
m sys} + \Delta S_{
m surr} < 0$  (nonspontaneous)

$$\Delta S_{
m univ} \equiv \Delta S_{
m sys} + \Delta S_{
m surr} < 0$$
 (nonspontane)  
 $\Delta S_{
m univ} = \Delta S_{
m sys} + \Delta S_{
m surr} = 0$  (reversible)

$$\Delta S_{\text{univ}} = \Delta S_{\text{sys}} + \Delta S_{\text{surr}} = 0$$
 (reversible)

### b./ Gibbs Free Energ

$$G = H_{\rm sys} - T \cdot S_{\rm sys} \text{ at constant T}$$

	$\Delta H$	$\Delta S$	$-T\Delta S$	$\Delta G$	Reaction Characteristics
ſ	-	+	-	-	at all T Spontaneous
١	+	-	+	+	at all T Nonspontaneous
	-	-	+	+or-	↓T Spon.; ↑T Nonspon.
١	+	+	-	+or-	↓T Nonspon.; ↑T Spon.

#### 7. Kinetics

#### 7.1 Reaction rate

Given general rxn:  $\alpha A + \beta B \longrightarrow \gamma C + \delta D$ 

$$0 < \mathsf{Rate} = \underbrace{-\frac{1}{\alpha}\frac{d[A]}{dt} = -\frac{1}{\beta}\frac{d[B]}{dt}}_{\mathsf{Rate of disappearance}} = \underbrace{\frac{1}{\gamma}\frac{d[C]}{dt} = \frac{1}{\delta}\frac{d[D]}{dt}}_{\mathsf{Rate of appearance}}$$

#### 7.2 Rate La

$$\boxed{ \text{Rate} \left[ \frac{M}{s} \right] = k[A]^m [B]^n }$$

- rate only depends on reactants
- k is the rate constant
- ullet m,n are the reaction orders
- m, n can be  $0, \frac{1}{2}, 1, 2, \dots$ • m + n is overall rxn order
- m, n are not necessarily equal to  $\alpha, \beta$

### '.3 Reaction Orde

Consider  $A \longrightarrow B$  $[A]_t = \text{concentration of } A \text{ at time } t$ 

$$[A]_t = \text{concentration of } A \text{ at time } t$$
  
 $[A]_0 = \text{concentration of } A \text{ at time } t = 0$ 

1st Order

$$\mathsf{Rate} = -\frac{d[A]}{dt} = k[A]^1$$
 
$$ln[A]_t = -kt + ln[A]_0$$
 
$$[A]_t = [A]_0 \cdot exp(-kt)$$
 2nd Order

2na Ora

$$\begin{aligned} \text{Rate} &= -\frac{d[A]}{dt} = k[A]^2 \\ &\frac{1}{[A]_t} = kt + \frac{1}{[A]_0} \end{aligned}$$



### Zero Order

Rate 
$$= -\frac{d[A]}{dt} = k[A]^0 = k$$

$$[A]_t = -kt + [A]_0$$

#### 7.4 Half Lif

Time needed for  $[A]_t = \frac{1}{2}[A]_0$ 

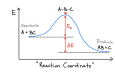
$$\underbrace{\frac{t_{\frac{1}{2}} = \frac{0.693}{k}}_{\text{1st Order}}}_{\text{1st Order}} \quad \underbrace{\frac{t_{\frac{1}{2}} = \frac{1}{k[A]_0}}_{\text{2nd Order}}}_{\text{2nd Order}} \quad \underbrace{\frac{t_{\frac{1}{2}} = \frac{[A]_0}{2k}}_{\text{Zero Order}}}_{\text{Zero Order}}$$

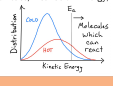
#### 7.5 Collision Mod

Reaction requires reactant molecules to collide with correct orientation and enough energy.

Higher  $T\colon$  reactants collide more often and with more kinetic E. Arrhenius

Molecules need minimum energy to react; Activation Energy,  $E_a$ 





#### 7.6 Arrhenius Equation

$$\begin{array}{l} \alpha A + \beta B + \gamma C \longrightarrow \text{products} \\ \mathsf{Rate} = k[A]^m[B]^n[C]^p \end{array}$$

$$k(T) = \underbrace{\left( \begin{array}{c} \text{collisions} \\ \text{per time} \end{array} \right) * \left( \begin{array}{c} \text{fraction of collisions} \\ \text{properly oriented} \end{array} \right)}_{A} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{with } E > E_{a} \\ \text{* } exp\left[ \frac{-E_{a}}{RT} \right] \end{array} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{with } E > E_{a} \\ \text{* } exp\left[ \frac{-E_{a}}{RT} \right] \end{array} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{with } E > E_{a} \\ \text{* } exp\left[ \frac{-E_{a}}{RT} \right] \end{array} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{with } E > E_{a} \\ \text{* } exp\left[ \frac{-E_{a}}{RT} \right] \end{array} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{with } E > E_{a} \\ \text{with } E > E_{a} \\ \text{* } exp\left[ \frac{-E_{a}}{RT} \right] \end{array} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{with } E > E_{a} \\ \text{* } exp\left[ \frac{-E_{a}}{RT} \right] \end{array} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{with } E > E_{a} \\ \text{* } exp\left[ \frac{-E_{a}}{RT} \right] \end{array} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{with } E > E_{a} \\ \text{* } exp\left[ \frac{-E_{a}}{RT} \right] \end{array} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{with } E > E_{a} \\ \text{* } exp\left[ \frac{-E_{a}}{RT} \right] \end{array} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{with } E > E_{a} \\ \text{* } exp\left[ \frac{-E_{a}}{RT} \right] \end{array} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{with } E > E_{a} \\ \text{* } exp\left[ \frac{-E_{a}}{RT} \right] \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{with } E > E_{a} \\ \text{* } exp\left[ \frac{-E_{a}}{RT} \right] \right]}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{fraction of molecules} \\ \text{fraction of molecules} \\ \text{fraction of molecules} \end{aligned} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{fraction of molecules} \\ \text{fraction of molecules} \end{aligned} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{fraction of molecules} \end{aligned} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{fraction of molecules} \end{aligned} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{fraction of molecules} \end{aligned} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{fraction of molecules} \end{aligned} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{fraction of molecules} \end{aligned} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{fraction of molecules} \end{aligned} \right)}_{*} * \underbrace{\left( \begin{array}{c} \text{fraction of molecules} \\ \text{fracti$$

A is a "frequency factor", assumed T-independent

$$ln(k) = \frac{-E_a}{R} \cdot \frac{1}{T} + ln(A)$$



#### 7.7 Reaction Mechanisms

A sequence of elementary rxns that sum to the overall rxn. The kinetics of elementary rxns are determined by how many molecules have to collide, referred to as the molecularity.

Elementary rxns have rate law where  $m,n,p\dots$  are equal to stoichiometric coefficients.

Molecularity	Elementary rxn	Rate law					
Unimolecular	$A \longrightarrow P$	k[A]					
Bimolecular	$A + A \longrightarrow P$	$k[A]^2$					
Bimolecular	$A + B \longrightarrow P$	k[A][B]					
tisten Reactions							

#### 1.8 Multistep Reaction

Overall rate law results from the individual rate laws for the individual elementary reactions.

$$\begin{array}{ccc}
2 & A & \longrightarrow & 1 + C \\
I + B & \stackrel{k_2}{\longrightarrow} & A + D
\end{array}$$
 elementary rxns
$$A + B & \longrightarrow & C + D$$
 overall rxn

Assume  $k_1 << k_2$ , i.e. step 1 is "rate limiting"  $\implies$  rate overall  $= k_1[A]^2$ 

#### 7.9 Catalysts

Substances that increase rxn rate, but are neither produced nor consumed in overall rxn.

Lower  $E_a$  has bigger impact  $\Rightarrow$  Appears in exponent of  $k(T) = A*exp(\frac{-Ea}{RT})$ 

#### 8. Chemical Equilibrium

$$A \xrightarrow[k_r]{k_f} B$$
 , at equilibrium: Rate  $= k_f[A] = k_r[B]$ 

#### 8.1 Law of mass action, equilibrium-constant

molarity concentrations partial pressures  $K_c = \frac{[C]^{\gamma}[D]^{\delta}}{[A]^{\alpha}[B]^{\beta}} \qquad K_p = \frac{[P_C]^{\gamma}[P_D]^{\delta}}{[P_A]^{\alpha}[P_B]^{\beta}} = K_c(RT)^{\Delta n}$   $K >> 1 \to \text{products dominate}, \ K << 1 \to \text{reactants dominate}$ 

 $K >> 1 \rightarrow$  products dominate,  $K << 1 \rightarrow$  reactants dominate K depends on T and is unitless **heterogeneous equilibria**: exclude pure solids / liquids from K reaction quotient Q **if not at equilibrium**, calculated like  $K_c$ 

- rxn written in reverse:  $K = K_{\text{original}}^{-1}$
- rxn multiplied by n:  $K = (K_{\text{original}})^n$ • multistep rxn:  $K = K_1 \cdot K_2 \cdot K_3 \dots$
- With catalysts: equilibrium is reached faster, K unchanged

#### 8.2 Le Châteliers principle

# Disturbance in concentration system reacts to consume added substance

system reacts to consume added substance Substance added



### Disturbance in pressure

reduced volume  $\rightarrow$  system shifts in direction with fewer moles of gas

# Disturbance in temperature Endothermic Exothermic increased T right shift left shift

 $\begin{array}{lll} \text{increased T} & \text{right shift} & \text{left shift} \\ \text{decreased T} & \text{left shift} & \text{right shift} \\ \end{array}$ 

### 8.3 Connection to Thermodynamics

$$\Delta G^{\circ} = -RT \cdot ln(K)$$
 or  $K = exp(\frac{-\Delta G^{\circ}}{RT})$ 

### 9. Acid Base Reactions

#### 9.1 Brownsted Lowry acids and bases

$$\underbrace{HA(aq)}_{\text{acid }(H^+\text{-donor})} + \underbrace{B(aq)}_{\text{base }(H^+\text{-acceptor})} \rightleftharpoons \underbrace{A^-(aq)}_{\text{conjugate base }} + \underbrace{HB^+(aq)}_{\text{conjugate acid}}$$
 strong acids/bases completely ionize, weak acids/bases don't

#### toionisation of water

Amphiprotic substances (ex. Water) can act as acid **and** base

$$\begin{split} \underbrace{H_2O(l)}_{\text{acid}} + \underbrace{H_2O(l)}_{\text{base}} &\rightleftharpoons \underbrace{OH^-(aq)}_{\text{conjugate base}} + \underbrace{H_3O^+(aq)}_{\text{conjugate acid}} \\ K_w &\equiv K_C = [OH^-][H_3O+] = 10^{-14}(25^\circ C) \end{split}$$

# 9.3 Reaction of acid with water / base with water acid-dissociation base-dissociation

$$K_a \equiv K_C = \frac{[A^-][H_3O^+]}{[HA]}$$
  $K_b \equiv K_C = \frac{[HB^+][OH^-]}{[B]}$ 

#### 9.4 p-Scales

$$p(\xi) = -\log(\xi)$$
  $pH = -\log[H_3O^+]$   $pOH = -\log[OH^-]$   $pH + pOH = 14$   $pH < 7 \rightarrow \text{acid}$   $pH > 7 \rightarrow \text{base}$ 

#### 5 common ion effect and Buffers

$$CH_3COOH + CH_3COONa \rightarrow \text{dissociates to } CH_3COO^ HA(aq) + OH^-(aq) \rightleftharpoons A^-(aq) + H_2O(l)$$
 $A^-(aq) + H_3O^+(aq) \rightleftharpoons HA(aq) + H_2O(l)$ 

$$[H_3O^+] = K_a \frac{[HA]}{[A^-]} \to pH_{\text{buffer}} = pK_a + \log\left(\frac{[base]}{[acid]}\right)$$

For disturbance small relative to  $[HA], [A^-] \rightarrow \text{small pH change}$ 

### 10. Redox Reactions

#### $10.1\,$ Oxidation Numbers

- Atoms in elemtal form: 0
- Monoatmic ions: ionic charge
- Nonmetals in ionic/molecular compounds: negative oxidation numbers

Oxygen: -2 (except peroxide ion,  $O_2^2$  -, -1) H: +1 (except if bonded to metal, -1)

F: -1 (always)

CI, Br, I: -1 (except if bonded to oxygen)

• Sum of oxidation numbers for atoms in compound equals its

 Sum of oxidation numbers for atoms in compound equals i net charge

### 10.2 Energy and Batteries

- Anode(-): where oxidation occurs
- ullet Cathode(+): where reduction occurs
- Anions migrate towards anode. Cations migrate towards cathode.

 ${\sf Electric\ potential} = {\sf potential\ energy\ difference\ per\ unit\ charge}$ 

$$1V = \frac{1.6 \cdot 10^{-} \, 19J}{1.6 \cdot 10^{-} \, 19C}$$

 $E_{\rm cell}^{\circ} \equiv$  Cell voltage at standard conditions  $E_{\rm cell}^{\circ} \equiv$  Potential energy available if reduced

transferred in balanced cell rxn

Cell potential  $\Rightarrow E_{\rm cell}^{\circ} = E_{\rm red}^{\circ}({\sf cathode}) - E_{\rm red}^{\circ}({\sf anode})$ 

### 0.3 Connection to Gibbs Free Energy

$$\begin{split} \Delta G^\circ &= \text{J/mol of rxn} \\ \Delta G^\circ &= -nFE_\mathsf{cell}^\circ \qquad F = \text{Faraday's constant} \\ &= 96'485\text{C/mol }e^-\text{'s} \\ E_\mathsf{cell}^\circ &> 0, \Delta G^\circ < 0 \\ \qquad n &= \text{unitless number moles of }e^-\text{'s} \end{split}$$

 $\Rightarrow$  Spontaneous!